DT05 Rec'd PCT/PTO 0 3 DEC 2004.

TRANSLATOR'S VERIFICATION

I hereby declare and state that I am knowledgeable of each of the German and English languages and that I and reviewed the attached made translation of International Patent Application PCT PCT/EP03/03448, filed on April 2, 2003, from the German language into the English language, and that I believe my attached translation to be accurate, correct to the best of my knowledge and ability.

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

November	18,	2004	D. Mullen	
Date			Signature	

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Method and agent for the prevention, inhibition and treatment of sepsis

The present invention relates to novel methods for sepsis prevention or sepsis treatment and the agents which can be used in such methods. It is based on a finding, namely that antibodies diagnostic novel (autoantibodies) which have the so-called properties of inhibiting "natural killer cells" (NK cells) were found with high sensitivity in the sera of patients suffering 10 from sepsis, and on the preventive and therapeutic measures which the Applicant has derived from the critical role which these antibodies can play in the development of the sepsis owing to their NK cellinhibiting properties. 15

In particular, the present invention relates to methods

for the prevention and treatment of a septic reaction in human patients (patients at risk of sepsis), which, after a medical intervention and/or a trauma (accident, burn, war injuries, decubitus and the like), a sepsis has developed or could still develop, and, 5 over and above this, also for avoiding a potential risk of a septic reaction in patients who have to undergo, for example, a surgical operation in which a sepsis must be feared as a dangerous complication, for example in the field of visceral surgery, transplantation 10 chemotherapy high-dose medicine and haematology/oncology.

The term "sepsis" is now used in close association with the term "inflammation". Inflammations are defined very 15 generally as certain physiological reactions of an organism to different types of external effects, such as, for example injuries, burns, allergens, infections by microorganisms, such as bacteria, fungi and viruses, to foreign tissues which trigger rejection reactions, 20 or to certain endogenous states of the body which in autoimmune trigger inflammation, for example Inflammations may occur and cancer. diseases harmless, localized reactions of the body but are also typical features of numerous serious chronic and acute 25 diseases of individual tissues, organs, organ parts and tissue parts.

Local inflammations are generally part of the healthy
immune response of the body to harmful effects, and
hence part of the life-preserving defence mechanism of
the organism. However, if inflammations are part of a
misdirected response of the body to certain endogenous

for example, in autoimmune ^ as, such processes, diseases, and/or are of a chronic nature, or if they reach systemic extents, as in the case of systemic inflammatory response syndrome (SIRS) or in a severe sepsis caused by infection, the physiological processes typical of inflammatory reactions go out of control and life-threatening actual, frequently the become pathological process. Regarding the modern definition of sepsis, reference may be made, for example, to the discussion of the definition of sepsis in K. Reinhart et al., "Sepsis und septischer Schock" [Sepsis and Intensivmedizin, Georg septic shock], in: Verlag, Stuttgart, New York, 2002, 756-760. According to this, "the clinical picture of sepsis is a separate pathophysiological and clinical entity which, also in terms of treatment, must be described and treated separately from the underlying infection".

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It is now known that the origin and the course of inflammatory processes are controlled by a cascade of substances which are predominantly of a protein or peptide nature or are accompanied by the occurrence of certain biomolecules for a more or less limited time. The endogenous substances involved in inflammatory reactions include in particular those which can be vasoactive cytokines, mediators, the assigned to hormonal and/or phase proteins acute substances, regulators. The inflammatory reaction is thus a complex which both endogenous reaction in physiological substances activating the inflammatory process (e.g. $\mathtt{TNF}-\alpha$, interleukin-1) and deactivating substances (e.g. interleukin-10) are involved.

In systemic inflammations, as in the case of sepsis or of septic shock, the inflammation-specific reaction cascades spread in an uncontrolled manner over the whole body and become life-threatening in the context of an immune response which is excessive or causes dysfunction.

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Whereas at least in Europe the systemic bacterial infection detectable by a positive blood culture long characterized the term sepsis, sepsis is now primarily understood as being systemic inflammation which is caused by infection but, as a pathological process, has considerable similarities with systemic inflammations which are triggered by other causes. Thus, the direct detection of bacterial pathogens was recently replaced or supplemented by complex monitoring of physiological parameters and also by the detection of certain endogenous substances shown to be involved in the sepsis process or in the inflammatory process, i.e. specific "biomarkers".

is which substance endogenous introduced An is as a sepsis biomarker particularly suitable procalcitonin. The determination of procalcitonin as a sepsis marker is the subject of the publication by procalcitonin al., "High serum et M. Assicot concentrations in patients with sepsis and infection", The Lancet, Vol. 341, No. 8844, 1993, 515-518; and the EP 0 656 121 B1 patents DE 42 27 454 C2 and and US 5,639,617.

The availability of the sepsis marker procalcitonin has given considerable impetus to recent sepsis research,

and intensive efforts are now being made to find the supplement biomarkers which can further of procalcitonin determination and/or are capable providing additional information for purposes of fine diagnosis or differential diagnosis. Results of these 5 efforts are to be found in numerous patent applications of the Applicant, in particular in DE 198 47 690 Al or WO 00/22439, and in a number of still unpublished Applications (DE 101 19 804.3 German Patent PCT/EP02/04219; DE 101 31 922.3; DE 101 30 985.6) 10 Applications (EP 01128848.7; European Patent EP 01128850.3; EP 01128851.1; EP 01128849.5; EP 01129121.8; EP 02008840.7 EP 01128852.9; EP 02008841.5). Reference is hereby made to the content patent applications said patents and 15 supplementing the present description.

substances during formed endogenous Since the inflammations are part of the complex reaction cascade of the body, not only are such substances of diagnostic interest but attempts are also currently being made, with considerable effort, to intervene therapeutically in the inflammatory process by influencing the origin and/or the concentration of individual substances of this type, in order to stop at as early a stage as possible the systemic spread of the inflammation which is observed, for example, in sepsis. In this context, endogenous substances which can be shown to be involved in the inflammatory process are also to be regarded as potential therapeutic targets. Attempts starting from certain mediators of the inflammatory process influence this therapeutically in a positive manner are described, for example, in E.A. Panacek, "Anti-TNF

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Anästhesie für Journal strategies", Intensivbehandlung; No. 2, 2001, 4-5; T. Calandra et al., "Protection from septic shock by neutralization of migration inhibitory factor", macrophage Medicine, Vol. 6, No. 2, 2000, 164-170; or K. Garber, 5 sepsis solution", may be "Protein С Biotechnology, Vol. 18, 2000, 917-918. In view of the fairly disappointing results of such therapeutic approaches to date, there is considerable interest in identifying further endogenous biomolecules which are 10 as inflammation- or sepsis-specific as possible and, as therapeutic targets, also open up new prospects for success for the prevention and treatment of sepsis.

15 A general object of the present invention can therefore be defined as providing novel measures (methods, agents, uses) for the prevention and treatment of sepsis, which arise from the discovery of biomolecules which occur with high sensitivity in sepsis and play a 20 key role in the development and in the course of sepsis.

This object is achieved by uses according to Claims 1 to 3 and embodiments thereof according to Claims 4 and 5 and agents according to Claim 6.

The present invention is based in general on the surprising experimental finding that a certain antibody or autoantibody type known per se in other contexts is found with high sensitivity and at significantly increased levels with extremely high frequency in sera of patients who had subsequently developed a sepsis or already had a sepsis, whereas the same antibody is not

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detectable or detectable only in substantially smaller amounts in healthy normal persons.

The fact that the antibodies found are those which can eliminate the function of "natural killer cells" (NK cells; cytotoxic lymphocytes) and that it has moreover been found that related antibodies determined in other scientific contexts cross-react with these antibodies which damage the NK cells plays a decisive role regarding the preventive and therapeutic measures described in this Application.

If such antibodies which inhibit NK cells are found at increased levels in virtually all patients suffering from sepsis, they are therefore a therapeutic target for the prevention of sepsis and treatment of sepsis. In the context of the present invention, as explained in more detail below, they can become the target of preventive, inhibitory and/or therapeutic measures with the use of measures, some of which are basically known agents according to the invention for per se, or fighting and binding pathogenic (auto)antibodies and rendering them harmless and/or of measures with the aim of specifically influencing the antibody formation by the immune system. Furthermore, their detection prior to a sepsis risk event may be a reason for taking greater safety measures, for example in the context of a preventive treatment with antibiotics, for avoiding development of a sepsis.

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More precisely, the present invention starts from the surprising results of measurements by the Applicant on sera of normal persons and patients suffering from

sepsis with the aid of a ligand binding assay with high sensitivity for antibodies binding to the gangliosides AG_{M1} and G_{M1} . The surprising result was that, in such measurements, antibodies binding to asialo- G_{M1} (anti- AG_{M1} antibodies) and/or antibodies cross-reacting therewith, in particular antibodies binding to monosialo- G_{M1} (anti- G_{M1} antibodies) of the IgG and/or IgA type were found substantially in all measured sepsis sera.

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The increased presence of antibodies which bind to asialo-G_{M1} and hence have properties which inhibit or damage NK cells in virtually all sera of patients suffering from sepsis indicates that such antibodies are directly involved in the development of a sepsis and/or can be correlated with an increased risk to a patient at risk of sepsis, i.e. with the risk that the patient will develop a sepsis or with its individual disposition to sepsis.

The physiological binding partners of said antibodies 20 are gangliosides. Gangliosides are glycolipids which are constituents of the extracellular side of the plasma membrane of animal cells and as such also occur in nerve tissue. They contain several monosaccharide units per mole but have no phosphorus content and are 25 assigned to the sphingolipids. Compared with proteins, they tend to be low molecular weight biomolecules. The gangliosides to which the antibodies discussed in the bind are the invention the present context of monosialo-ganglioside referred to generally as G_{M1} and 30 in particular the associated "asialo" compound AG_{M1} . G_{M1} has a polysaccharide chain of 4 sugar monomer units units, one comprise two D-galactose which

N-acetylgalactosamine unit, and one D-glucose unit, the latter being bound to a so-called ceramide moiety. In the ganglioside G_{M1} , an N-acetylneuraminic acid radical (NANA; sialic acid or o-sialinic acid radical; "monosialo" radical), which is missing in the sialinic acid-free asialo- G_{M1} (AG_{M1}), is bound to the D-galactose unit arranged inside the polysaccharide chain.

Said gangliosides and related compounds are associated with numerous important biological functions of the human body, including, for example, axonal growth and neuronal differentiation, receptor functions and participations in various immune reactions of the body and in signal transduction and cell-cell recognition.

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antibodies or that known been Ιt long autoantibodies which bind to the ganglioside G_{M1} , in structures, and to particular its carbohydrate which structures of other molecules carbohydrate resemble these ("simulate these") can be detected in the human body in certain contexts. The physiological role of such antibodies and their possible importance for clinical diagnosis are the subject of numerous scientific investigations. However, the detection of anti- G_{M1} antibodies has not to date been correlated with disadvantageous physiological reactions which important for the development of sepsis and which may attributable to properties, known per οf se, antibodies binding to asialo-Gm1.

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By far the predominant part of all published papers are concerned with the role and the diagnostic significance of anti-ganglioside antibodies in neuropathies, for

example in immunomediated motor neuropathies, such as (radiculoneuritis, syndrome Guillain-Barré (Miller-)Fisher related the and polyradiculitis) occurrence of anti-Gm1 increased An syndrome. autoantibodies in some patients was also reported in 5 association with Alzheimer's disease. Furthermore, they were found in individual HIV patients. That they also occur with very high sensitivity in cancer is a still unpublished novel finding of the Applicant, on which inventions disclosed in European the 10 Applications EP 02009884.4 and EP 02009882.8 are based. Reference is hereby made to the two last-mentioned European Patent Applications and the list of references for EP 02009884.4 for supplementing the statements in the present Application. 15

The findings which form the basis of the inventions in the present Application, and the preventive and therapeutic measures derived therefrom, are explained in more detail below.

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In order to avoid unjustifiably narrow and restrictive interpretations of the terms used in the present Application and the associated claims, some of the most important terms are to be defined in particular below for the purposes of the present Application:

without includes, term This "Antibody": distinguishing between different methods of genesis and formation, antibodies 30 external antigens and both against structures, against endogenous autoantibodies, where the latter may

also have become autoantibodies by antigen cross-reactions from antibodies against external antigens and may have preserved their binding capability with respect to external antigens.

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When, for example, it is stated that an ganglioside binds "to antibody and to antigen structures structures simulating ganglioside structures" or is gangliosides "reactive towards certain gangliosides", where reactive means "reactive in the context binding", it should specific sufficiently defined by this definition example, its specific without, for binding also to additional other antigen practical its structures, or (for reagents determination using immobilization marking or as orcompetitors) with molecular structures which only simulate AGM1, in particular the carbohydrate structure thereof, playing a role for the definition as antibodies according to the invention.

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"Ganglioside" In the context of the present invention, the term "ganglioside" primarily represents the gangliosides AGM1 in the characterization of the binding behaviour of the antibodies to be determined. However, the term is also intended to include related gangliosides

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not investigated to date, if it is found that antibodies also binding to these gangliosides and having a comparable diagnostic significance are found in sepsis sera.

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"Simulation"

in the context of the present When, stated that, in it is Application, addition to the specific gangliosides (AG_{M1} and G_{M1}), substances or compounds properties "simulating" binding effects thereof may also be used, what is that there are is meant by this carbohydrate compounds which have structures (including bacterial toxins) and which bind to the antibodies in question like said gangliosides. compounds can therefore be potentially used, like the gangliosides themselves, for specific binding of the antibodies (for example for the purpose of removing them from the blood circulation) or for their blocking (and hence rendering them

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substances context, stated In "simulating" the binding behaviour of gangliosides can be found, for example, with the aid of a screening method in which a biological sample (in particular high antia serum) which has ganglioside antibody titre and whose binding behaviour was determined in a

harmless).

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given assay is brought into contact with a candidate substance to be tested and the antibody binding is determined again assay. A substantial same reduction or elimination of the antibody binding in the assay is an indication that the substance investigated is "ganglioside-simulating substance". This carried out in also be can test association with establishing a patent infringement by agents which correspond specifically disclosed those claimed in the present Application.

- 15 Further meanings of the terms are evident to a person skilled in the art from the introductory and following description of the invention and its embodiments and the literature cited therein.
- In the description below, for explaining the diagnostic findings which are the basis of the preventive and therapeutically useful teachings, reference is made to figures which show the following:
- of the results of the graph Fig. 1 shows 25 measurement of antibodies of the IgG class which bind to monosialo-G_{M1}, in sera of 137 control persons, compared with the results of sera of patients the measurement of 89 suffering from sepsis; 30
 - Fig. 2 shows the results of a measurement of the same sera as in Fig. 1 for antibodies of the

IgA class which bind to monosialo-Gmi;

Fig. 3 shows the results of the determination of antibodies of the IgG class which bind to asialo-Gm1, in sera of 30 normal persons (controls), compared with the results of the measurement of 20 sera of patients suffering from sepsis (all sera are partial groups of the sera measured in Figures 1 and 2);

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- Fig. 4 shows the results of the determination of antibodies of the IgA class which bind to asialo- G_{Ml} , in the same sera as in Fig. 3.
- The invention is based on measurements of sera carried 15 out by the Applicant, substantially only the principle and the results obtained being measurement reproduced in the present Application. A detailed disclosure of the measurements carried out is to be prior unpublished European Patent the 20 found in Application 02009884.4 of the Applicant and a further application of the Applicant filed simultaneously with the present Application. Reference is made to both applications for supplementing the disclosure of the present Application: 25
 - 1. Determination of (auto)antibodies of the IgG and IgA type which bind to the gangliosides AG_{M1} and G_{M1} in sera of healthy normal persons (controls) and patients suffering from sepsis.

Using test tubes coated with gangliosides (AG_{M1} and G_{M1}) (GA-CTs), the free binding sites of which had been

saturated with BSA, series measurements were carried out on control sera and test sera. As described in more detail in the prior unpublished Application of the Applicant EP 02009884.4, and a further European Patent Application of the Applicant filed simultaneously with the present Application, antibodies from the respective serum (control serum and test serum) were bound, in a first incubation step, to the gangliosides used for the for separate determination coating, and then, the bound type, IqA and IgG the antibodies of antibodies were detected using marked animal (goat) anti-human IgG or anti-human IgA immunoglobulins. The detected immunoglobulins were marked bound quantified (in relative units, based on serum having the highest concentration of the respective antibody to of their basis the determined) on (acridinium ester as chemiluminescence marker).

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In order to obtain a measured signal representative of the amount of antibody, it is necessary to subtract a background signal which was determined under identical measuring conditions for the respective identical serum using those test tubes which, apart from the lacking ganglioside coating, were identical to the test tubes used for the actual measurements.

137 control sera (blood donor sera and - for avoiding age-related influences on the antibody concentrations - sera of normal persons of different ages from old people's homes and of the Applicant's employees) served as control sera for the antibody assays using GA-CTs which were coated with G_{M1} . For the antibody assays using GA-CTs using GA-CTs which were coated with AG_{M1} , a partial

group of these sera which comprised only 30 sera was measured.

89 sera of patients suffering from sepsis served as 5 test sera for the antibody assays using GA-CTs which were coated with G_{M1}. Exact clinical documentation existed for each test serum. For the antibody assays using GA-CTs which were coated with AGm1, a partial group of these sera which comprised only 20 of the sera of the patients suffering from sepsis was measured.

The results of the determinations of antibodies of the IgG and IgA classes using GA-CTs which were coated with G_{M1} are shown by way of example in Figures 1 and 2.

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The corresponding results obtained using GA-CTs which were coated with AGm1 are shown in Figures 3 and 4.

It should be expressly pointed out that numerous sera 20 in which significantly increased anti-AGm1 or anti-Gm1 antibody titres were found originated from patients from whom the blood sample had been taken shortly (about 2 h) after the sepsis risk event and who developed the typical sepsis symptoms only later on.

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should furthermore be pointed out that, experiment determine also to the corresponding antibodies of the IgM type analogously determinations of the antibodies of the IgG and IgA type, no levels for such antibodies of the IgM type which were increased to diagnostically relevant a extent were found in the sepsis sera (results not shown).

2. Discussion of the findings and measures according to the invention for sepsis prevention and treatment

impressively shown by the measured summarized in Figures 1 to 4, the determination of antibodies of the classes IgA and/or IgG which bind to gangliosides (AG_{M1}) and/or G_{M1}) permits distinction of the control group from the patients suffering from sepsis, by virtue of the fact that substantially increased AG_{M1} and G_{M1} antibody titres are found in virtually all (Fig. 1: 82 out of 89, i.e. 92%; Fig. 2: 80 out of 89, 90%; Fig. 3: 19 out of 20, 95%; Fig. 4: all 20, 100%) of the sepsis sera investigated.

15 The detection of substantially increased concentrations of antibodies of the IgA and IgG type also in patient's serum samples which had been obtained only a short time 2 h) after the "sepsis risk event" operation, accident, burn), and the lack of evidence of 20 antibodies of the IgM type, ruled out the possibility that the detected antibodies were formed only as a result of the "sepsis risk event" or of a bacterial infection associated therewith. However, this means that either the antibodies were already present 25 beforehand in the respective patient suffering from sepsis and/or that the activation of the presensitized immune system of the patient in the manner of a "booster" effect, triggered by the sepsis risk event, led to intensive antibody production.

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The fact that the AG_{M1} or G_{M1} antibodies were found at substantially increased levels in substantially all measured sepsis sera (90 to 100%) is therefore to be

interpreted so as to mean that the formation of a sepsis is either due directly to the presence of the antibodies in question in the respective patient or is at least the consequence of activation of the "molecular machinery" (in the form of B-cells) already present in the patient owing to prior immunization, which begins its intensive antibody production under the influence of the "sepsis risk event" or of an infection associated therewith. Patients without these antibodies or the presensitization required for their rapid production probably do not develop a sepsis or develop one only with difficulty, based on the experimental results to date.

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15 The observed results can be explained: it is known that so-called "natural killer cells" (NK-cells: cytotoxically active lymphocytes) have, on asialo-G_{M1} structures surface, to which antibodies can specifically bind and thus deactivate 20 and destroy the NK-cells. This is even routinely used the field of animal experiments which employs experimental animals in which tumours are to artificially produced to eliminate the immune defence of the experimental animal by administering anti-AGm1 25 antibodies in combination with a carcinogen or a tumour nucleus, so that the experimental cancer - desired in the animal model - can develop (Hugh F. Pross et al., Role of Natural Killer Cells in Cancer, Nat Immun 1993; 12:279-292: Lewis L. Lanier et al., Arousal 30 inhibition of human NK Cells, Immunological Reviews 155:145-154; Yoichi Fuji 1997, Vol. et al., Antibodies to AsialoGM1 Are More Sensitive than IgM Antibodies to Kill in vivo Natural Killer Cells and

Prematured Cytotoxic T Lymphocytes of Mouse Spleen, Microbiol. Immunol. Vol. 34(6), 533-542, 1990; N. Saijo et al., Analysis of Metastatic Spread and Growth of Tumor Cells in Mice with Depressed Natural Killer 5 Activity by Anti-asialo GM1 Antibody or Anticancer Agents, J Cancer Res Clin Oncol (1984) 107: 157-163; Sonoku HABU et al., Role of Natural Killer Cells against Tumor growth in Nude Mice - A Brief Review, Tokai J Exp Clin Med., Vol. 8, No. 5, 6: 465-468, 1983; Lewis L. Lanier, NK Cell Receptors, Annu. Rev. Immunol. 10 1998, 16: 359-93; Theresa L. Whiteside et al., The role of natural killer cells in immune surveillance of cancer; Current Opinion in Immunology 1995, 7:704-710; Tuomo Timonen et al, Natural killer cell-target cell interactions, Current Opinion in Cell Biology 1997, 15 9:667-673).

However, active NK-cells play an extremely important role in the human immune defence, also in the case of 20 severe bacterial infections. sepsis or Thus. example, Shuiui Seki et al., in: Role of Liver NK Cells and Peritoneal Macrophages in Gamma Interferon and Interleukin-10 Production in Experimental Bacterial Peritonitis in Mice, Infection and Immunity, Vol. 66, No. 11, 1998, 5286-5294, describe the important role of 25 NK-cells for the production of inflammation-promoting anti-inflammatory cytokines. They show that switching off NK cells artificially the with experimental use anti-AG_{M1} antibodies of 30 inhibition of the production of the anti-inflammatory interferon-y. Effects of surgical stress endotoxin-induced sepsis on the NK-cell activity have already been investigated, in particular in: P. Toft et al., in: The effect of surgical stress and endotoxin-induced sepsis on the NK-cell activity, distribution and pulmonary clearance of YAC-1 and melanoma cells, APMIS 1999; 107:359-364. A possible influence of physiologically formed antibodies with NK-cell reactivity, however, is not taken into account in any of the papers mentioned.

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The scientific literature does not reveal any discovery 10 which might suggest the method according to invention. Ιt is true that a determination anti-ganglioside antibodies in connection with severe acute infectious diseases was described in some papers. Such an infectious disease is Chagas disease caused by 15 the parasite Trypanosoma cruzi (cf. D.H. Bronia et al., in: Ganglioside treatment of acute Trypanosoma cruzi infection in mice promotes long-term survival and parasitological cure, Annals of Tropical Medicine & Parasitology, Vol. 93, No. 4, 341-350 (1999) and the 20 literature cited therein). The last-mentioned paper speculates that an observed, substantially advantageous effect of administering exogenous ganglioside to mice infected with the parasite T. cruzi might induce in said mice production of anti-ganglioside antibodies, 25 which then react with glycolipids of the membrane T. cruzi and should thus result in the death of the parasite. Owing to the findings in the present Application, such an explanation is not very probable: owing to their NK cell-inhibiting effect, the anti-30 ganglioside antibodies observed in the case of Chagas disease are not, as assumed, a healing-promoting factor instead a disease-inducing or disease-promoting factor. As a result of the administration of exogenous

gangliosides, which as such cannot be regarded as antigens, the anti-AG_{M1} antibodies are in fact not formed but probably blocked. Consequently, the effect of the NK cells may be restored in the mice (or in patients), and the immune system can overcome the parasite. All papers with which we are familiar establish no relationship at all between anti-asialo- G_{M1} antibodies and the origin and worsening of sepsis and therefore can neither anticipate nor suggest the method according to the invention.

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This also applies to other papers which show that administration of exogenous gangliosides or certain ganglioside derivatives in experimental animal models 15 anti-inflammatory effect orsignificantly increases the survival rate of mice to which the bacterial toxin LPS had been administered (cf. Silvia G. Correa et al., in: Anti-inflammatory effect of gangliosides in the rat hindpaw edema test, European 20 Pharmacology, 199 Journal of (1991)93-98; Amico-Roxas M., et al., in: Anti-inflammatory action of AGF44, a ganglioside ester derivative, Drugs Exptl. Clin. Res. XVIII(6) 251-259 (1992); James J. Mond et al., in: Inhibition of LPS-Mediated Cell Activation In 25 Vitro and In Vivo by Gangliosides, Circulatory Shock 44:57-62 (1995). In none of the papers are the observed effects of the interactions of the administered gangliosides with anti-ganglioside antibodies explained. The latter are not even discussed in the 30 given context.

Such an interaction in the context of antibody blocking can conclusively explain the findings described. The

conceptual principles of the present invention can be intensified as follows: it is to be assumed that the patients who are suffering from sepsis and whose sera were measured have the specific antibodies predisposition for their rapid production (in 5 the context of sensitization) before the "sepsis risk event". The sensitization of a patient with respect to the production of anti-ganglioside antibodies may have taken place, for example, as a reaction to some exogenous, antigenic stimulus (for example a general 10 infection with Campylobacter jejuni or Heliobacter pylori or, if appropriate, corresponding environmental substances), independently of the subsequent sepsis. It may remain latent for a long time thereafter. However, 15 once the production of anti-asialo-Gm1 antibodies has initiated or increases greatly in been individual, this patient has, as a disposition, preconditions that there will be damage to the NK cells and hence the immune defence in certain physiological stress situations, such as infections and other events 20 with high NK cell activity (for example cell degeneration through mutagenic events; a sepsis risk situation), by the anti-AGm1 antibodies and antibodies cross-reacting therewith. There is an increased risk that a defence reaction which is triggered by, for 25 "sepsis risk stress" and example, intervention by the NK cells will also stimulate the production of the above-mentioned antibodies, and that these will then cancel out the effect of the NK cells. The regulatory cycle of the immune response is then 30 decisively disturbed, and a sepsis may develop.

The detection of naturally occurring anti-AG $_{ t M1}$

antibodies and anti-ganglioside antibodies reacting therewith, e.g. anti-Gm1 antibodies, and the increased levels of such antibodies in sera of patients therefore mean suffering from sepsis that antibodies may represent a previously unconsidered parameter influencing the infection- or inflammationspecific cytokine cascade, in that they intervene in cytokine regulation cycle and, the natural disturbing or switching off the NK-cells, can cause this to malfunction and trigger a septic reaction in the patient.

It is therefore to be assumed that the anti-AG_{M1} or anti-G_{M1} antibody titres found at significantly increased levels in all sepsis sera which were measured in the above-mentioned assays constitute one of the preconditions for the origin of a sepsis, and the presence of such antibodies has a sepsis-inducing or sepsis-enhancing effect.

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The determination of such antibodies is therefore suitable for determining the risk situation (individual disposition) and for the prognosis in a patient who is to be classed as a sepsis risk patient.

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A method which comprises determining the presence and/or the amount of anti-AG $_{M1}$ antibodies and antibodies cross-reacting therewith in a biological fluid of a sepsis risk patient, and taking sepsis-preventing measures in the event of their detection, is derived therefrom. Thus, for example, a preventive treatment with antibiotics can be carried out for rapid preventive therapeutic intervention.

An administration of suitable substances for said purposes can also be carried out as a purely routine preventive measure, i.e. without prior determination of the antibodies. However, this is not very advisable in view of the seriousness of an incorrectly estimated risk of sepsis.

Since, without external stress, the antibody titres may in certain circumstances be very low, it is within the scope of the invention to carry out the antibody determination after an in vivo stimulation of the antibody formation of a patient at risk of sepsis, for example before a surgical operation, using safe stimulants. In view of the IgA antibodies found at substantially increased levels (cf. Figures 2 and 4), the antibody determination should also be capable of being carried out expressly by suitable assays in body secretions (e.g. saliva, mucous).

20 The increased titres of anti-ganglioside (auto)antibodies in patients suffering from sepsis are therefore of serious medical importance, namely as a pathogenic factor. With this knowledge, it is possible to develop novel, promising therapeutic approaches:

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1. Measures for the intracorporeal blocking of the pathogenic antibodies by administering to the patient agents (which of course must have the required compatibility) which are suitable for saturating ("for blocking") the antibodies binding to AG_{M1} .

The gangliosides themselves are such agents, and their good compatibility for human patients is known from

other contexts in which they have already been administered (they are administered in large amounts to patients suffering from Parkinson's disease). Where are said to such gangliosides have already been administered in relevant contexts for therapeutic purposes to humans without the above-mentioned contexts having been recognized, the agents used are to be excluded from the scope of protection of the present invention.

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In addition to the administration of the gangliosides themselves, in dosage forms and amounts suitable for the purpose according to the invention, ganglioside—"simulating" substances can also be administered, as already explained above. One possibility for identifying such substances, in particular those which bind more strongly to the antibodies than the gangliosides themselves, is described further above, in association with the definition of "simulation".

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Candidates for such substances may be, for example, oligosaccharides which consist of the sugar tetramer of the AG_{M1} molecule - without ceramide and sialinic acid radical - or contain these, and derivatives thereof.

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2. Removal of the antibodies from the patient's circulation. This can be effected, for example, by extracorporeal binding to solid affinity materials of ("plasmapheresis"). For the production plasmapheresis affinity columns used as binders, it is also possible, if appropriate, to use high-affinity carbohydrate structures which bind the antibodies with higher affinity than the gangliosides themselves and,

owing to their toxin properties, cannot be administered to a patient. Regarding the question of the production of T-cell anergy, reference may additionally be made to the content of EP 705 107 A1 and the further references mentioned therein.

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3. Reduction of the endogenous production of specific antibodies by administration of agents blocking antigen-presenting cells or for producing These measures may be accompanying 10 T-cell anergy. effects of the measures mentioned under 1., but may advantageously also be applied in a targeted manner by choosing other administration routes (e.g. oral) amounts which are suitable for the desired reduction of the antibody production without being sufficient for blocking all antibodies present in the circulation of a patient at risk of sepsis or a patient suffering from sepsis.